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Attachment A

Lise-Lotte Gundersen, *Tetrahedron Letters*, Vo. 35, No. 19, pp. 3155-3158, 1994 6-Chloropuines and Organostannanes in Palladium Catalyzed Cross Coupling Reactions

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6-Chloropurines and Organostunnanes in Palladium Catalyzed Cross Coupling Reactions

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Abstract: Carbon-circles band formation in the purios 6-position can easily be accomplished by pulsations enalysed creats coupling increases 6-chieveparties and organizations without protection of the poster sing Aby quantum. This backetique provides a convenient norm to passet cytokinhous.

Rinetin (6-durfaryhandropoulos) and roland 6-substituted purious, called cytokinine, and known to stimulate call growth and cell division in plants. I Several purious with potent cytokinin activity countin a 6-sibaryl or 6-sibyl miteriousts. An efficient name to these compounds would be transloss named camply and cross compling of organometallic magnets with readily available 6-haloporines. For instance, 6-chloroporine la (Scheme 1) is communicity available and the substance can also satily be prepared on a large scale. Because of the acidity of the purion ring NH function (pEa of 6-chloroporine; 7,50),4 organometallic magnets with low basisty are desired. Examples of practices meet metal mediated couplings of purious with a free NH function are restricted to palladium catalyzed Heck coupling of terminal alkynes with haloporines, and scircle entalyzed coupling of methylthioporine with Originard reagents in which case 2.5 equivalents of the basic Grigosod inagents were required, 5 Palladium catalyzed oross coupling between heteropyl halides and organotic derivatives is an established surfaced for carbon-carbon bond formation to heterocycles, 7 but the scope of this method in surely hydrotigated in the purious ring pyracm, 8 Only coupling structions employing N-9 alkylated independents are reported. In fact, very recently the first two examples of coupling in the purious appeared, 8

Palladium camiyzed cross compling of organosmumanes Zu - a with 6-chicopymines In - b has been studied (Scheme 1, Table 1).

Scheme 1

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Table 1. Pd-Cambyzeri Coupling hotteren Organizationness and Chloroparium.

	Purme I	Organostaname 2	React time (b)	Tenn. (C)	Product 3	Yield (%)
	là	PACH=CHSubba ₃ 2n ¹⁷	20	QS,	₩	34
0	. 3b	2	45	50	H _N N N N N N N N N N N N N N N N N N N	75
	12	CELECTOR()States 24	20	80	* The *	<i>3</i> 8
	Xa.	PhSoBu ₃ 2c	20	100	M 3 3 4	81
	ль	2c	45.	100	H _N N N N N N N N N N N N N N N N N N N	73
@	1a	(2-12:00y)5aB-3 24 ¹²	20	90	\$\frac{1}{2}\dagger*	81
	ъ	21	45	90	H ₂ N N N N N N	76
	Ja.	PhCH ₂ SaBu ₃ 2n ¹⁵	20	130	N N M	36

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Organostaments are known to tolerate a wide variety of functional groups on the coupling partners, and the stammarks 2n - a coupled with the relatively acidic parises 1n and 1h. Bis(triphenylphosphine)palladium(II) dichloride was found to be a suitable catalyse, 10

This methodology makes purious with alternyle, anyle, and hencestyl substituents in the 6-position castly accessible, and the potent synthinise 6-rans-styrytestine 3s, was prepared in a one-step procedure superior to the methods reported outlier. 10 Byon the much less reported being (hibstyl) stansane 2s participated in the combine to give the 6-beary/spanie 3h in a moderate yield.

The 6-myrel perions 3a • b were prepared from a 1 : 9 minimum of cis and cross etyroids respect 2a (1.3 equiv.), but only the cross products were formed as judged by ¹H NMR of the cross products, indicating a slightly higher reactivity of the cross organicis respect. Isomerization of the alternal in the seasion can not be excluded, but resention of the alternyistments double bond geometry is penetrally observed in pallacitum catalyzed orom couplings.⁹

The order of reactivity of anyl halides in palladium mentioned cross coupling reactions is found to be Ar-I > Ar-Br >> Ar-CL? Only anyl chlorides with an electron deficient theo carbon will participate in the reaction. For instance, 2- and 4-chloropyrimidiates are shown to be sufficiently activated for the coupling to boom,? In ractive reports, independent have been amployed in couplings with organosimments, and even though the yields in Heek couplings of 6-chloroporines are nather moderate compared to the corresponding independents, 3.13 the present study shows that the 6-chloroporines As and 1b are reactive example to give high yields of the coupling produces in palladium catalyzed reactions with allocayle and anylin derivatives. This is a great advantage state todoprehas are less available. In fact, independent are often prepared from the corresponding chloroporines 15.16

The lower reason rate observed in the couplings of 2-amino-5-chloroporine 1b compared to 6-chloropurine 1a might be anothered to higher electron density in the former ring, making the compound least activated for medeophilic attack from Pd(0). Initially, the coupling mentions with the 2-amino compound 1b were carried out in a much loss concentration one to poor sulmbility, but mentions of the parise 1a acoust not no be significantly affected by concentration.

In Summary, carbon-carbon bond formation in the puriou 6-position can easily be accomplished by pallacinus cambyard cross coupling of readily available 6-chloroperines with organostances without protection of the puriou ring NH function. This technique provides an officient route to potent cytokinines.

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10. The organizationness: 2 (1.3 mmel) was added to a mixture of bis(triphosyl dishloride (39 mg, 0.00 ramol) and 6-chloroportee 1a (1.0 morel) in dry DMF (6 ml) or 2-amino-6chloropurhes 1h (1.0 mmol) in DMF (15 ml) and the mixture was sourced under N2 at the temperature indicated in Table 1, before the reaction mixture was filtered and evaporated, and the residue was washed with other. The captle product was purified by Hash chromatography or receystallization. The Ly-NMR (DMSO-de) 5 6.32 (s. TH), 7.4 - 7.5 (m. 4H), 7.7 (m. 2H), 2.10 (s. 1H), 8.25 (d. / 16 Hz. 1H), 12.6 (be, 1H). MS (BL): 237 (2 M*), 207 (100). 3c: 1H NIMR (DMSO-dg) & 1.46 (LJ 7 Hz, 3H), 4.15 (g, J 7 FL, 2H), 4.77 (4, J 2 Hz, 1H), 5.62 (4, J 2 Hz, 1H), 8.77 (5, 1H), 8.91 (4, 1H), 13.6 (64, 1H), MS (EL); 190 (100, M²). 3m ¹H NMB (DMGO-de) 5 6.58 (s, 2H), 7.5 - 7.6 (m, 3H), 8,11 (s, 1H), 8.7 - 8.5 (m, 219), 12.7 (bs, 125). MS (ELL): 211 (100, M*). 3£ 1H NMR (DMSO-da) 8 7.36 (m, 1H), 7.92 (m, 1H), 8,66 (m. 227), 8,84 (s, 181), 13.7 (hs, 183), 148 (B.L.): 202 (100, M²). 3g: ¹87 NHCR (DMSC)-4&) 5 6,40 (s. 2H), 7.28 (sc. 1H), 7.20 (sc. 1H), 2.12 (s. 1H), 2.57 (sc. 1H), 12.7 (bc. 1H), 163 (E.1): 217 (100, 167), Labordia, J. W.; Seile, J. K. J. Aus. Chem. Soc. 1923, 103, 6129-6137.

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